

) PATENT ABSTRACT (11) Decument No., AU-A-47099/89) AUSTRALIAN PATENT OFFIC

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COSMETIC OR PHARMACEUTICAL COMPOSITION CONTAINING MICROSPERES OF POLYMERS OR OF FATTY SUBSTANCES FILLED WITH AT LEAST ONE ACTIVE PRODUCT

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Claim.

cosmeric or pharmaceutical composition containing such microspheres illed with active product(s) in a suitable carrier can be employed for ringing medications to a determined point of the body, in particular for polication to the skin. However, topical application does not generally eve the desired effectiveness because the epidermis forms a barrier.

scording to the present invention, it has been found that, if the crospheres of the cosmetic or pharmaceutical composition are chosen from particular size range, the effectiveness of the active product which they intain is greatly increased in a very unexpected manner. Studies nducted by the Applicant Company have made it possible to establish that is considerable improvement was linked with the entry of the microspheres to sebaceous follicles.

im 1. Pharmaceutical or cosmetic composition for topical application staining, in a suitable carrier, microspheres of polymers or of fatty setamces with a melting point higher than 50°C filled with at least one live product, characterised in that at least 80 % by weight of the respheres have a diameter of between 3 µm and 10 µm.

(11) 47098/89

- 2. Composition according to Claim 1, characterised in that the polymer is chosen from the group consisting of styrene-based polymers, -alanime-based polymers, polymers derived from acrylic or methacrylic acid, polymersms derived from lactic and/or glycolic acid, crosslinked proteins and proteins accepulated by heat.
- 5. Composition according to Claim 1, characterised in that the fatty substance is chosen from the group consisting of fatty alcohole and derivatives of alcohole and of fatty acids.

Form 10

COMMONWEALTH OF AUSTRALIA PATENTS ACT 1032-60

COMPLETE SPECIFICATION

(ORIGINAL)

	Class	Int. Class
Application Number		
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Complete Specification Lodged:		
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Complete Specification for the invention entitled:		

The following statement is a full description of this invention, including the best method of performing it known to be used to the statement of the following statement is a full description of this invention, including the best method of performing it known to be used to the statement of the following statement is a full description of this invention, including the best method of performing it known to be used to the following statement is a full description of this invention, including the best method of performing it known to be used to the following statement is a full description of this invention, including the best method of performing it known to be used to the full description of this invention.

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COSMETIC OR PHARMACEUTICAL COMPOSITION CONTAINING MICROSPHERES OF FOLIMERS OR OF FATTY SUBSTANCES FILLED WITH AT LEAST ONE ACTIVE PRODUCT

COSMETIC OR PHARMACIUTICAL COMPOSITION COMPARTISMS MICROSPHERES OF POLINERS OR OF FACET SUBSTANCES PINIOR HITH AT LEAST ONE ACTIVE PRODUCT.

The present invention relates to a cosmetic or pharmacoutical composition containing microspharus of polymers or of fatty substances filled with at least one active product in a suitable carrier.

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It is known in the otate of the art to prepare microcapsules in which the active principle is enclosed and is not in contact with the external environment (see particularly French Patent 2,219,036 and Chiropean Patant 315,054). However, at the time of application, the microcapsule can break prematurely and release the active principle immediately.

It is also known to propare natural or symthetic polymers in the form of microspheres by crosslinking these polymers in numpension. A process for the manufacture of poly-3-alamine microspheres is described, for example, in Franch Palant 3,530,350. It is also known to prepare microsphares of fatty substances.

It is also known that these microspheres are capable of filling with chemical products, in particular with active products (see particularly tha abovementioned Franch Patent and US Patent 4,590,825). In the present application, an active product means any product having an activity from the cosmetic or pharmacoutical visupoint. The colld product forming the soluble. This carrier can be an aquecus solution or an

oily phasa.

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A commetic or phermacoutical composition containing such discrepances filled with active product(s) in a suitable carrier can be amployed for bringing medications to a determined point of the body, in particular for application to the skin. Eccepter, topical application does not generally have the desired

effectiveness because the epidermia forms a barrier.

According to the present invention, it has been found that, if the nicrospheres of the counstic or pharmacounical composition are chosen from a particular size range, the offectiveness of the active product which they contain is greatly increased in a very unexpected manner. Studies conducted by the Applicant Company have made it possible to establish that this considerable improvement was linked with the entry of the nicrospheres into sebaceous follicies.

The subject of the present invention is therefore a cosmetic or pharmacontical composition for topical application containing, in a suitable carrier, microspheres of natural or synthetic polymers or of

Satty substances with a malbing point higher than 80°C, filled with at least one active product, characterized in that at least 90 % by unight of the dispendence amployed have a dispense of between 3 µm and 13 µm.

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In fact, microsphores which have a diamet of a minpange defined above enter the sebaccous follicle, but little into the skin. The said microsphores, sherefore, selectively and progressively meach the follicular canal, where the active product which they carry diffuses into the follicular canal and the surrounding cisaues. On the other hand, the substract forming the microsphore is subsequently rejected by virtue of the flow of sobum and/or of the growth of heir. Any undestrable reaction of the organism towards the solid compound forming the microsphores is thus avoided.

It should be noted that, when the nicrospheres have a dissetor smaller than 3 µm, they also enter the follicular canals, but the horny layer as well, it a high concentration. Now, this release of the active principle in the horny layer, for our ple in the case of antiacne preparations, is reflected by the appearance of secondary effects which are undestrable insefar as the active product is released in the application and which surround the follicular channels; whereas, in the case of medications as ing systemically, the active product is released in a conveniently the active product is released in a conveniently, the active product is released in a conveniently, the active product is released in a

barrier intervenes. Owneall, therefore, in both cases, the release of the active principle in the horry layer corresponds to a reduction in the effectiveness of the composition. Then the microspheres have a diameter greater than approximately 10 µm, they remain restly localized on the surfaces of the skin victous intering it, resulting in an imaginativeness of the topical application, since the active product can only be released on the horsy layer. In both cases, the targeting of the active products is markedly inforier to that which is obtained by making use of the

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In other words, the invention proposes to select the size of the microapheans do as to promote their selective entry into the sebaceous follicias; in the case of acce, the active product is thus brought specifically to the target regions without undesirable secondary effects on the healthy skin regions supposeding the follicular channels; in the case where the active product is a medication which lots systemically, the follicular channel constitutes a highly efficient route of general administration insofar as the diffusion of the active product into this compartment emerges onto a highly vascularized region.

It was not obvious that dicrospheres capable of satering the heir follicle had to have the dimensions defined above. In fact, the new distance of the

pilosabaccous orificas is included in a size range which is quite different from that indicated above in the case of the microspheres; for example, on the forehead, this average diameter is between 52 µm and 32 ma. In man, the surface area of the pilosebeceous 5 orifices situated on the forehead is approximately 0.002 am2 (N.J. Cunliffe, W.D.H. Perera, P. Thackray, M. Williams, R.A. Forstor and S.M. Williams, British Journal of Dermatology, 1976, 25, 133). Assuming that the contour of the follicular channel is approximately 10 circular, the average diameter of the pilosebaceous crificas can be astimated, according to this paper, at 50.5 µm. This diameter, redetarmined by the Applicant Company by measurement of the size of the pilosebaceous orifices situated on the skin of the forehead of six 13 healthy volunteers, is found to be between 52 μm and 92 µm (see study described in test 3 of the present application). This considerable difference between the range of the diameters of pilosebaceous orifices and 12 the range of diameters of the effective microspheres made the invention particularly surprising for the specialist. This surprising nature is furthermore confirmed by the fact that in the abovementioned US Patent 4,690,825, the size indications supplied are 25 simed only at microspheres which have diameters of between 10 and 100 pm.

The microspheres which have the desired size can be selected by ecroaning, especially in a moist medium,

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microspheres obtained by a process giving microspheres which have a more extended sange of sixes. It is also possible to obtain microspheres whose sixes are contained in the desired range by suitably directing the process for the manufacture of the microspheres. The sixe of the microspheres can, for example, to adjusted by choosing the polymerisation solvent and the crosslinking agent, or by modifying the rate and the time of stirring of the reaction medium. These various modifications form part of the state of the art and/or are within the competence of the specialist.

The natural or synthatic polymers which can be employed for the manufacture of the microspharus of the composition of the present invention are chosen from those capable of being applied to the skin without undesirable effect and cape to of forming microspheres which have the desired dimensions. They must also be compatible with the active product employed.

The polymers which can be employed in the compositions of the present invention may be adventageously chosen from:

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- styrane-based polymers, such as polystyrane;
- \$-alanine-based polymers, such as poly-\$-alanine;
- polymers derived from acrylic or methacrylic acid:
- polyesters derived from lactic and/or glycolic acid;

- proteins crosslinked:

either by glutaraldehyde or by an acid dichloride such as tarephthaloyl chloride,

or in the presence of an activator such as a carbodiimide;

- proteins coagulated by heat (albumin).

The polymers which can be employed are preferably chosen from polymers based on poly- β -alanino and polyesters derived from lactic or glycolic acid.

The fatty substances which can be employed may be chosen from:

- derivatives of alcohols and of fatty acids, such as tristearin, semisynthetic triglycerides or glycarol monostearate;
- 15 fatty alcohols such as cetyl alcohol.

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The fatty substances which can be employed are preferably chosen from fatty substances which have a melting point of approximately between 50°C and 100°C.

The active products which can be employed in the composition according to the invention are those liable to be applied to the skin. They may be chosen from:

- agents for treating acne, such as compounds with action of retinoid type (vitamin A, retinoic acid or its derivatives);
 - bensoyl peroxide;
- growth factors of paptidic nature, such as the proteinic or epidemic growth factor (EGF):
 - skin-reinforcing agents, such as bensyl

nicotinata;

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- agents for treating hair, in particular antiloss or hair regrowth agents, such as minoxidil and antiseborrhooics such as S-maxboxymethyleys bind or octopizon;
 - antifungals such as nystatin or aconasole;
 - Astringents, such as aluminium chloride;
- antibiotics, such as enythromycin and tetracycline;
 - antivirals, such as vidarabina;
- antihypertensors, such as clonidine hydrochloride;
 - m antianginals, such as nitroglycerias;
 - vasodilators, such as bradikymin;
- agents for treating cardiovascular disorders, such as peptides of the tachykinins group, for example 'substance P';
- Antiinflarmatory agents, such as aspirin or hydrocortisons and its derivatives;
 - antiallergens such as chromoglycatos;
- antiprurities, such as phenothiesine
- derivatives;
- neurostimulants, such as caffgine or theophylline;
- antidepressant agents, such as lithium salts and, more perticularly, lithium carbonats;
- natural compounds employed in n urobiological research, such as capsaicine;

- ansesthatics, such as lidocains and procains;
- hormone staroids such as 17-a-ocatradiol and 17-\$-osatradiol.

The suitable carrier is in aqueous form or in the form of oil.

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The carrier in aqueous form may be an aqueous gelchtsined with the aid of a gelling agent, such as the crosslinked polyacrylic acid sold under the trade name "Carbopol" by Goodrich BF or the cellulose derivatives sold under the trade name "Mucel" by Enroules; or a hydroalcoholic gel containing, for example, propylene glycol. It is also possible to use a lipophilic aqueous solution such as an aqueous solution of silicones.

The cils which can be employed as carriers are liquid or scaleolid cils such as triglycerides of C_0-C_{10} fatty acids and their mixtures, vaseline, liquid paraffin and lanolin.

The pH of the carrier is preferably adjusted to a basic value.

The carrier is in the form of liquid, of gel, of cream, of pants, of ponade or of dry powder. To obtain a pasts, a pomade or an ointment, an excipient is added, such as polyethylene glycol, a war such as beeswax or lanolin.

The commetic or pharmacoutical compositions according to the present invention generally contain from 1.3 to 40 % by weight of microspheros, at loant 80 % of which have dismeters of between 3 and 10 pm.

They also contain from 0.03 % to 40 % by weight of active product.

The microspheres are manufactured by any known process. The polystyrene microspheres are widely marketed. Those of poly-#-alanine can, for example, be prepared according to the processes described in French Patent 3,530,350.

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To introduce the active product into the microsphere, the active product is dissolved in a solvent or a mixture of solvents which have a sufficient affinity for the compount forming the microspheres. Among the suitable solvents, especially for poly-s-alanine apheres, there may be mantioned, for example, water, glycarol, ethanol, disthylene glycol, acetone and, in general, reter-miscible organic solvents.

When a solvent has been employed to obtain the microspheres filled with active product, the said microspheres may be employed as such or after removal of the solvent remaining therein. This polyent may have remained therein as solvent of the active product and/or as a swelling agent for the microsphere itself when the polymer of which it is made is liable to swell in the said solvent. When the microspheres are employed after removal of the solvent, the active product remains nevertheless trapped in (or on) the microsphere on drying. Swelling of the polymer by a solvent produces microspheres in gel form, provided that the

quantity of solvent does not exceed certain limits, which are different depending on the polymer of which the microspheres are made. The microspheres filled with at least one active product, be they dried or not, are mixed with the chosen certain.

The cosmetic or pharmacoutical composition obtained is applied in the usual way to the skin, preferably with a gentle massage. In an alternative form, the microspharms are filled with an active product in ionised form: in this case, after application of the composition to the skin, the release of the active product may be accelerated by ionophoresis.

The examples given below, purely by way of illustration, no limitation being implied, will allow the invention to be better understood. Tests A, B and C are measurements provided to explain the remarkable effectiveness of the compounds according to the invention.

Tost A:

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In this test, the size of the pilosebaceous orifices in man is evaluated. This study was carried out on six healthy volunteers (three men and three women) aged from 25 to 35 years, and it was carried out on the skin of the forehead.

After having carefully cleaned with soap a region of skin of approximately 2 cm², a dye (dark brown direct dye "L'Oreal removative", marketed by the company known

as 'l'Oreal') is chosen and is applied, for Sifteen minutes, to the left or right side part of each subject's forehead. At the end of the exposure time, the coloured region is cleaned with a little water to remove the excess dye. This region is photographed with a macrophotographic assembly produced with the aid of an Olympus owners. This apparatus makes it possible to take standardized photographs of the region to be analysed (same distance and same magnification for all the subjects). The dye employed is no longer wisible 24 hours after the application.

The distribution of sizes of the pilosebaceous orifices is established by image analysis with the aid of the 'Quantimet 520' apparatus from Cambridge Instruments, from transparencies of the forehead. The apparatus measures the surface area S of the follicle openings and calculates the diameter D of each follicle according to the formula:

D - 2 (S/s)

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30 The results and given in Table 1 below.

The average diameter of the follicles is found to be between 52 μm and 82 μm for all the subjects studied.

Tast B:

rests were carried out to establish the relationship between the siss of the microspheres and their entry through the borny layer and the follicles of the human skin.

Thase tests employed fluorescent polystyrene microspheres of various calibres between 1 μ m and 24 μ m which had the characteristics given in Table II below. These batches of polystyrene microspheres were suspended at a concentration of 10 3 by weight in 3 mixture of triglycerides of C_4 - C_{10} fatty acids marketed under the trade mark thygliol 912° by Dynamit Mosel) the tests were performed on the face lift skin of the face of famile patients aged from 44 to 65 years.

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SURLEUT RO.	254	SERVE	CIDACTER (pd)		
		ralias	'mazzd.		
			+/- attendend	<90 3	<95 %
			<u>ರೋಚಿಸುವಾ</u>		
					1
1	Ħ.	105	R2 +/- 34	<128	<150
3	3	102	63 +/- 42	<120	<141
3	P .	111	32 +/- 43	<133	<158
6	7	115	52 ÷/- 25	< 87	< 99
5	Ħ	103	79 +/- 37	<129	<143
6	H	68	79 +/- 31	<124	<132

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	(هند)		(۵۵سر)	Dre.	escauca
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	1.17	1	40.0	17458	pricht pina
10	3.1	3	0.1	17155	Telles-green
	6.83	7	0.2	18161	Astron-Assess
	7.0	7	0.3	17156	yallow-groon
	9.13	9	0.6	13140	yello-gran
	9.55	10	1.53	18142	yello-, men
. <u>Te</u>	23.8	24	4.3	15241	respondant

- The sime consequence of these particle size standards were complied by (61 (Polyacianos Dr.).
- ee Phinnescame type: (see Table III).

TABLE III

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Pluoroscanca	Excitation max.	Emission sax.
Bright blue	365	453
Yellow-green	458	540

The applications are carried out, approximately & hours efter surgical excision, on facial shin which was not been deep-fromen (atorage at 4°C in a cold chamber). The skin is freed from its subcutaneous tissue with a scalpel, and is then slightly attratched 5 and pinned onto a support cover d with aluminium. The cutaneous surface is carefully cleaned by wiping with a paper handkerchist, followed by a slight 'stripping' carried out with the adhesive tope sold under the trade name "Transpore". The various suspensions of microbeads 10 are then applied with a glass spatule for 15 minutes, with 5 minutes' massage, inside 2.5-cm2 application sites delimited by plastic rings bonded using a cyanoacrylata polymer-based adhesive marketed under the name of "Cyanolit". At the end of the application time, 1.5 the emcass product which has not entered the skin is removed with a cotton-stick followed by three wary slight applications, to the surface of the skin, of a piece of adhesive tape of trude name "Transpore" (adharing little to the skin and causing no 10 delamination of the horny layer). Biopsies of the application sites, as well as of a control skin region without application, are taken with a "Punch biopsy" punch 6 mm in diameter and are frozen in liquid nitrogen. The entry of the microbeads into the horny 25 layer and the follicles is then demonstrated, using a fluorascanca optical microscopa (photomicroscopa IIIRS, Seiss, Hear Germany) on deep-frozen vertical skin

sections 10 µm to 15 µm in thickness, produced with the aid of a crycalcrotome (Cryostat Bright, Bright Instrument Company Limited).

The results obtained are the following:

- microspheres 24 µm in diameter remain localised on the surface of the skin without entering it;
- microspheres 9 µm to 10 µm in diameter have a tendency to collect around the follicular canals;
- 7-mm microspheres have been able to be selectively placed inside the dabaceous follicles;
- microsphares from 1 µm to 3 µm have a tendency to enter both the horny layer and the follicles.

Test C:

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Tests were carried out to establish the relationship between the size of the microspheres and their entry into the horny layer and the follicles in the rat.

These tests employed poly-j-alanine microspheres; three samples which had average dimmeters of approximately 2 μm , 5 μm and 12 μm respectively were tested.

The microspheres employed are prepared by crosslinking poly-s-alanine with the sid of glutaraldehyde. This synthesis is described in French Patent 2,530,250. These microspheres are then made fluorescent by an intermediate reaction of hexamethylenediamine with the residual aldehyde fur tional groups present at their surface, followed by

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a reaction with dansyl chlorids. The microsphares obtained exhibit a very homogeneous powerful green fluorescence in ultraviolet light. These microsphares have the following characteristics:

- sample l: dismetor = 1.79 ± 0.95 مر (50 % balow (2.9);

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- sample 2: diameter = 4.8 : 1.1 pm (90 % below 5.1 pm); prepared according to Example 1;
- sample 3: diameter = 12.4 ± 2.2 µm (90 è below 15.1 µm).

These size measurements were determined by the fluoresence image analysis technique using a "Quantimet 520" apparatus marketed by Cambridge Instruments Co.

The application protocol employed is the following: after anaesthesia with pentoberbital (30 mg/kg dose), 5-cm² application sites are delimited by a plastic ring bonded adhesively to the back of the ICO female nude rat (170-180 g average weight). The various suspections are applied for 2 hours, in a quantity of 5 to 10 mg/cm², inside these sites. In order to tast the influence of r sage on the antry of the poly-s-alanine microspheres into the sebaceous follicles, the application is carried out by comparing two massage periods: one minute and five minutes. The entart is bound throughout the experimental period in order to avoid any contact with the region of application. At the end of 2 hours, the excess product which has not entered the skin is carefully removed

with a cotton-stick; three very slight applications of a gisca of adhasive tape of trade name 'Transpore' (adhasing litely to the skin and causing no delamination of the horny layer) to the skin surface are then carried out. Singuies of the application regions are taken (6 mm in diameter) and from in liquid nitrogen. The entry of the microspheres into the horny and follicular compartments is then established using the fluorescence optical microscope on deep-from vertical skin sections from 10 mm to 15 mm in thickness, produced using the cryomicrotoms.

In order to test the influence of the carrier on the entry of the poly-s-malanine microspheres, the latter are formulated, at a concentration of 10 % by weight, in the following carriers:

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1) Acueous gal which has the following
formulation:
Crosslinked polyacrylic acid sold under the trade
name "Carbopol 940" by Goodrich BF 0.4 g
Sodium hydroxids (aqueous solution at a
concentration of 10 % by weight) 2.0 g
Water q.s. 100.0 g
2) Water-silicone carrier consisting of:
Water 5.0 g
Silicone oil sold by Dow Corning under the
moderance "Q2-3225c" q.s. 100.0 g

a) in succession in the aqueous gel, microspheres

2 µm in diameter enter the various layers of the horny layer as well as inside the following canals. S-µm microspheres are rarely present in the larny layer, after one minute's manage, and are located rather at the entry of the following canals; this tempency to enter the following is slightly more prenounced after S minutes' massage. Microspheres 12 µm in diameter enter neither the horny layer nor the following can is.

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b) the water-silicons carrier has an influence on the entry of the 2-µs microspheres; the latter are more numerous inside the sebectous follicles and exhibit a uniform distribution in the horny layer. On the other hand, with this derrier, practically no 3-µs microspheres are found in the horny layer; they are located very deep in the follicles in the vicinity of the sebectous glands; in this case, massage also has a beneficial influence on the entry of the microspheres into the follicular compartment. As in the case of the aqueous gel, microspheres 12 µs in diameter enter neither the horny layer nor the follicles.

Examples 1 to 5 below describe processes for the manufacture of poly-\$\rho\$-alanine microspheres, fly rescent or filled with active products and having the desired diameter.

Example 1

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Franciation of fluorascent coly-a-alaning microaphoras Stance 1: Preparation of poly-a-alaning aphoras in auspension.

1125 g of toluena, 444 g of tert-butanol and 0.75 g of corolymar (octadocens/salsic anhydrids) (sold under the trade name 'PA-18' by Gulf) are introduced into a 3-litra radctor equipped with an anchor-type stirrar with a diameter of 90 mm, a nitrogen inlat, a dropping funnal and a distillation column head. After heating this mixture to 70°C, 180 g of acrylamide are added. The temperature is then raised to 100°C and 90 ml of the assotrope mixture (water/toluone/tortbutanol) are distilled off. After the end of distillation, the reaction mixture is cooled to 30°C and the stirring rate is adjusted to 600 reg/min. A solution of 3.30 g of potassium text-butylate in 62 g of tart-butanol is then added ever 10 minutes. The dropping funnal is minused with 75 g of tolugna. After stirring for 5 hours at 80°C, the material is allowed to roturn to ambient temperature. 11.25 al of concentrated hydrochloric acid are than added dropwise to the mixture.

Stage 7 : Croselinking of the poly-s-alanine apheros.

42 g of an aqueous solution containing 25 % of luteraldehydo are odded to the suspinsion of poly-8-alanda alcrospheres thus obtained, ove 30 minutes, with stirring at 600 rev/sin and at a temperature of

30°C. After animing has been continued for 4 hours at this temperature, the suspension is allowed to return to ambient temperature.

After settling, the supermatant solvents are removed and the microspheres are vashed twice with 300-ml portions of ethanol. Draining after each washing is carried out by contribuging at 3,500 mev/min. A vashing with 15 litres of water is then carried out continuously and the water is then removed to a final mixture volume of 600 ml is reached.

The crosslinked poly-A-slamins is then dried by freeze-drying and 135 g of a white powder are obtained, in which the diameter of the microspheres is on average 4.80 t 1.1 µm, determined by the image analysis technique using a "Quantimet 520" apparatus markoted by Cambridge Instruments Co..

Stage C : Reaction with 1,6-diaminchers a

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20 g of 1,6-disminohamme are added to a sus insion of 20 g of the poly-g-alamine spherus obtained in stage B in 100 g of water. Stirring is continued for 24 hours at ambient temperature and the material is then drained on a no. 4 glass sinters lastly, it is washed with water until the aqueous washers are at a neutral pH.

Stage D : Fixing of the fluorescent product.

The microspheras obtained in stage C are suspended

in 80 ml of ps 8.9 buffer solution (270 ml of 0.1 K

massCo. solution brought to pH = 3.9 by adding

approximately 30 ml of 0.1 H solution of Halch). By of dansyl chloride in solution in 80 g of acetons are introduced into this suspension. The Mixture is heated for 10 minutes at solvent reflux and is then drained on a no. 4 glass sinter and finally washed with acetone until all traces of dansyl chloride have disappeared from the solvent wash, monitored by DV datection at 250 nm. The opheres are first dried in air and then under meduced pursuance as ambient temperature. The final colour or the microspheres is light yellow.

Elample 1

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Propagation of colu-1-alemina microsobares filled with bancovi narowida

Signal: Preparation of poly-d-alanine aphores in suspansion.

This stage is identical with stage A of Example 1. <u>Stage B</u> : Orosalinking of the poly-s-alumina microspheros

of glutaraldabyde are added standily over 13 minutes to a suspension of poly-\$\beta\$-alanine microsphere: obtained in stage \$A\$, kept vigorously stirred (\$600 rev/ain) and at a temperature of \$0°C. After stir ing has been continued for \$4 hours at this temperature, the suspension is allowed to return t embient temperature. After settling, the supermittent episonts are removed and the microspheres are washed twich such stahing is carminious.

by centrifuging (3,500 ray/min). Mashing with 13 littee of water is then carried out continuously and the water is then removed until a final mixture volume of 600 ml is reached. The swellen polymer is finally dried by freeze-drying and 132 g of white powder are obtained, in which the diametal of the microspheres is on average 4.05 i 2.02 µm, measured according to the same method in Stage 3 of Example 1.

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Stags C : Reduction of the mesidual aldehyde functional groups.

2.2 litres of water are added to 150 g of crosslinked poly-s-alanian microspheres obtained in stage 3 and are homogenised by stirring. After cooling to a temperature of between 5 and 10°C, a cooled solution of sodium borohydride in water (5.2 g of Mans, in 600 ml of water cooled to 8°C) is added slowly. The reaction mixture is kept between 5 and 10°C for 5 hour and the p3 is then brought to 7 by adding scenic acid.

After contribuging the sixture and dispersing the solid residue in 450 ml of water, it is subjected to continuous washing with 5 litres of water (washing in an "Amicon" cell equipped with a 0.2-pm Diapor filter, pressure 2 bars, stirring throughout the washing). The hydrated microspheres are then divided by firesee-drying. The absence of colour in the presence of Schiff's reagent makes it possible to semulude that the residual aldemyde functional groups have been reduced. After analysis, the diameter of the sicrospheres is identical

with that of the original microspheres.

Stage D : Introduction of the active product.

44.5 g of benseyl peroxide (75 % by weight grade) are dissolved in a minture made up of 1125 g of accesses and of 375 g of water, 30 g of the microspheres prepared in stage C are then suspended in this solution. The suspension is concentrated in a rotary evaporator at reduced pressure, at a temperature not exceeding 35°C, to a total weight of 262 g of suspension.

The benzoyl peroxide content of the suspension obtained is 9.1% by weight.

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Pretaration of poly-1-alemina misroapherea filled with benevi misotimeta.

Stages A to C : Preparation of the microspheres.

Stages A to C are carried out as in Iz sple 2.

Stage D: Introduction of the active product.

2 g of beneyl nicotinate are dissolved in a mirture made up of 40 g of water and 40 g of ethanol; 10 g of microspheres prepared in Stage C are then suspended in this solution. The suspension is kept stirred for 2 hours and the otherol is then removed in a rotary evaporator, the temperature being maintained at a value bolow 35°C. Finally, the microspheres are dried by fromse-drying.

Transland
Francisco of roland-daning microsphores filled with

banayl nicoticata.

Stages A to C are carried out as in Example 1 and stage D for introducing beneat nicotinate as active product, as in Example 3.

Enspela 3

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Extracte acid

Stages A to C : Preparation of the microspheres.

Stages A to C are carried out as in Example 3. <u>Stage D</u> : Introduction of the active product.

15 mg of butylhidroxytoluens (antioxidant) are dissolved in 30 g of 1,2-propylens glycol at a temperature of 30°C. 24 mg of rotinoic acid are dissolved in 10 g of the mixture obtained above, at ambient temperature, under argen and in the absence of light. The solution obtained in filtered with the mid of 0.2-pm "Millipore" filteres. 3 g of the microspheres graphed in stage C are suspended in this solution in

His is carried out with a spatula. After two hours' absolution, a yellow powder is obtained. Detarmin tion of retinoic acid in the spectrophotometer ($\lambda = 353.8$ nm) after description of the active principle into directly sulphoxid.

the absence of light and under a stream of arcon.

Theoretical concentration : 0.15 %.
Calculated concentration : 0.157 %.

The gol to drough with optiming and them drough-dried.

Calculated monomorphism 11.3 % (by UV

determination of 330 an after suspending in otherwip.

Sacrata 3:

5 274PARTETION of their subsessed microsophurus filled mich medicale again.

<u>deach</u>): Propulation of the solution of active principle.

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300 mg of all-trans methods acid ar, dissolved in 3 ml of 1,3-dichiprosthame, in the absence of light.

Stage 3: Coating of the active principle with fatty substance microspheres.

4.75 % of triatuaria and 250 mg of glycorol monostograta are introduced into a stainless steel masetor provided with a mitrogen inlet and equipped with a magnatic stirror and a heating plate. Mixing is carried out by scirsing at a temperature of 30°C. The solution of active principle prepared in stage A is thin idead in the absence of light. The mirture obtained is large stirred at 50°C and is than blown, under a nitrogen pressure of 7 base, into a spraying monals connected to the reactor Reparatus *1/4 JCC-55-3U.B153-55', Zaani). The microspheres commisting of the retinoic acid coating with the minture of tristearin-glycarol conceteerate fatty substances are than domaid downwards of this spraying possis inside a filtration chamber (longth: 05 cm) and are than collected on a grid (Millipore, 34 on in diameter,

preferably, "1 TT30 293 001). A pollow-coloured possess is obtained. The rotinois acid contant of the microspheres obtained is 2.78 % by smight. The disaster of the microspheres, determined by image analysis (1937-Videoplan apparatus, Montron) is 4.43 % 1.30 ms.

Complete to 13 balow relate to the proparation of despected or pharmacourisal compositions from Microspheres filled with active product and propared in Examples 3 to 3.

naphtheir hold

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3.3

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0.3 g or poly(lactide-co-glycolide) sold by
Dupont under the trade size of 'Nodicorb sold bi' and
5 mg of 6-(0-(1-adamanty)-4-methoxyphonyl))-2-naphthoid
acid are dissolved in 13 ml of methylene chloride. The
exquite solution obtained is emploified with mechanical
attirting (3000 sov/min) in 100 ml of an aquacus gel
containing 0.3 g of hydroxypropyl collulose sold by
Aqualon under the trade name of "Riccel SF". Deckanical
extirting is continued for 3 hours, which permits the
progressive and complete symporation of methylene
chlors s.

The microspheres obtained are recovered, essied times since with distilled rater and dronze-dried, the size distribution of the microspheres obtained by this perhod in analysed with a distribution.

Trample :

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Exercision of colverability of a restrict that with along the before leading

To my 1 to 7 : Proparation of the dicrospheres.

Stages A and I are carried out as in Emanyly 3. Stages 3 : Independentian of the action product.

37.6 no of elemidiae hydrochloride are discolated in 13 g of water in the absence of light, and 3 g of microspheres propared in stars C are then added to 12 g of the above solution. Mining in carried out with a spately. After 2 hours, absorption a white powder is obtained. The microspheres are then dried by freezeddrying. Determination of eleminical hydrochloride in the finished product is carried out by EDEC analysis after description of the active principle.

Calculated concentration : 1 %.

73 3731 71

Propartition of poly-feelings pleroupherne dilled with mich

20 <u>Gramme 1 to ff</u> : Proparetion of the microspheres.

Stages A and C are carried out as in Stample 1. Stage D: Introduction of the active product.

2 g of minoxidil are dissolved at 30°C in a minure made up of 75 g of athanol and 75 g of mater.
3 g of poly-#-minutes apheres obtained according to swaps C are interded into this solution. The minture is arthred for 1 hour in the notary evaporator and the solvent is then evaporated eff until a gel is obtained.

the aphares is between 1 and 13 pm, with he promage disc of 5 pm; core than 10 % of the microsykares baye & discourse of heteres 3 and 10 pm.

The encapsulation is checked in the lobbering manager

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- 1) improvious of the democratical principle, the accepted of the expects of society principle,
- 3) inapocaine by alectron aderescopy confirms the theanes of organals autoids the spheres and the absence of organals on the surface of the spheres.

To avaluate the doques of accessulation of the active principle is the microspheres, a sample of the microspheres obtained above (100 mg) is extracted with settlebydrosuman (8 ml); is is then filtered; the dilinate is analyzed by high performance liquid chromatography; the degree of analyzedation of 5-(3-(1-3demancy)-3-machomyphenyi)]-2-amphibits acid in 6.71).

Transla 10 :

Translation of mainth price-co-demolical minmaphorus

filled with regions acid

Microspheres filled with retinoic acid can be obtained by the same method of preparation as in themple 0: the 5 mg of 5-(3-(1-adamanty1-4-methomychangl))-3-maphthoic acid and then replaced by 3 . 4 of vertinoic acid.

Cornela II 1

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3.5

Proposation of reinformation discountered dilled with the Improved the new Language of the Control of the Contr

Diomonyhous are propared from triglyconicos, massly a systematest palm oil marketed under the name of "Mostican abla" by Americ Dobbi, by A systeming proposes with the mid of a pressurited appropring unic.

The interpretation and the metry principle, namely in-banaylphonylacesemperatoride at a denomination of 13 % by weight relative to the seight of integly various, are maltim at 30°C under sittingen atmosphere and in the absence of light in a thermostated stai issuested absence. The molton mixture is propelled with mixtures (0.3×10° MRs pressure) up to the aposts at a contain also rate and the appropriaging current dear at the absolution mixture alternative cut at the absolute under mitrogen pressure (0x10° MRs pressure).

The openying is considered in a sealed stainless steal messel which has a temperature gradient from approximately =150°C at the bottom to 10°C at the top. This gradient is created by provious interest stion of liquid microgen into the bottom of the wessel.

has a general rule, depending on the type of notale which is chosen, the apraying altregen pressure and the slow rate of the liquid determine the average dismeter of the apheres obtained. Thus, the lower the slow rate, the smalless the droplets leaving the notale and, denses pointly, a microspheres as the bottom of the wages! The thompson, the higher the spraying pressure,

the smaller the diameter of the openies and the torn home-mesons the ciss electrobustion.

In this energie, uniform microspheres are obtained without tree express; of active principle which are vicible made the interescent. The distrator of the made waring from 1 to 15 ps, with a use, distrator below 10 pm. The properties of active principle uncompensed, decorated by high performace liquid phase shromatography, was established to 13 d.

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11. 护室 医肝炎

.. god is propared by mixing the following increminates:

Manmpla 11

A gol is propared by mining the following ingradients:

catara completely, ' to daily new 10 days, this
proparation has one the Authors proparation.
Company of the compan
I get is prepared by mining the following
is a stable to the stable to t
Moreophore to rangion propared according to
<u> </u>
Crossiinked polyscrylie sold sold undan the funcu
amme "Curbonol 940" by Goodries 37 28 6
Mabbs gud 4.5 km
ವರದವುದು ದಿನದಾರವಾಗಿದ್ದಾರೆ. ನೀಡು ಕಾರ್ಯಕ್ಕೆ ಕಾರ್ಯಕ್ಕೆ ಬಿಡುವುದು ಬಿಡುವುದು ಬಿಡುವುದು ಬಿಡುವುದು ಬಿಡುವುದು ಬಿಡುವುದು ಬಿಡುವ
then opplied to the other by ceasuage matil it
enters occupiately, twice daily for 00 days, this
prinparation has appoliant applicant groupartion.
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A got to propaged by utiling the following
ingrociants:
Microspheros propared according co
Emmode 3 (as camyas are mecosasty) & J
Groupliniad polyacrylic acid pold under the trace
name "Carroy of 980" by goodrich ar 9.4 g
Hator q.b 580 g
Sodium hydroxida q.3 93 = 7
When applied by massage match it interme
completely, trice daily for 30 days, to cortain parts

of the body, for example the broasts, this proparation

contributes to usking this limes.

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	When applied to the which by researce motal to
	estars completely, water daily der 30 days, Mil:
	programmion has ormaliant metiaena proparties.
	The same of the sa
2.	a gel so grandend by mining the rollering
	ingrodiants
	- Tulerouphaman ebrained in Grampia : 10 ;
	- Colletosa corivativas sold ancher the trace name
	naledal by Bermine
()	- Natur g.s
	Wear applied to the ski by massage matil it
	octars completely, brice daily for 2 to 3 words, this
	proparation has expellent autilypertensive proporties.
	The second of th
. 3	A gol is propaged by mining the following
	ingrediante:
	- Microspheres obtained in Mample 7 17.24 7
	a, Colimbone domivations sold mader who exact mana
	"Elucal" by Harculus 1.88 g
:-5	- Tatar 13.32 g
	≥ Empgylana glycol q.2 188 g
	This gal is applied tories daily so a scalp which
	and andergone a considerable hair loss. Ester 3 menute.
	undatment at a sate of 1 ol per application 1
: 5	significant improvement is moted.

1222-11-22

	a gal de impeared by manie; the following
	Lagrodionts:
	a Microspineou constinta in Unimpla 1 1.3 m
3	" Triffelona gazinasites noid mozam tos amedo moza
	"Mucal" by Marville (
	- Mater g.s 120 q
	Those applied to the own by manager with the
	emplers armadistrily, remon emply done so days, their
70	proparation has accellent mileton proparties.
	<u> 1895212-1</u> 3
	a gal is propared by mining the collecting
	ingrodiants:
	= literesphanes propared secondition to
3 .5	35x5pla 315 g
	- Crosslinkod polysergiic sold sold moder the name
	. Carbopol 940° by Goodrich 37 6.4 q
	- Retar 4.3 200 g
	- Jodina kydromido q.J gs = ?
13	When applied to the skin by massage antil in
	enters completely, twice daily for 30 days, thin
	properation has excellent entisons properties.
	<u> </u>
	a gel is prepared by mixing its sollowing
15	ingrodients:
	- Granoppeans bandured recompant to
	notable 19 a 9
	- Crossitained polyacrylic wid sold under the same

	"Carbopol 940" by Geodrien 37	3.4 g
	- Bator G.J	100 g
	- Josim Agaranian g.s	74 a 1
	Thou applied to the older by names or tackle i	t
5	onther samplestin, which dully for 36 days, which	
	propagation has accultant ancasens propagation.	
	<u> </u>	
	n gel he propered by thining was following	
	ingrodiants:	
10	- Microsphures proping Jecorating to	
	<u> </u>	20 g
	- Crosslinked polyacrylic acid sold under the	numa
	"Carbopol 340" by Goodwith M	0.5 g
	교 위송원5호 (제·점· ·································	100 9
15	- Sodium hydromida q.s	pE = 7.
	When applied to the akin by massage ascil i	.T
	carars complessly, water tally for 30 days, this	ı
	promotion are explicate serifative recommendate account	æiad.

the group consisting or present for treating law, a principality agants, agants for treating law, artification, asking and artification, asking agants are interpretable to a proper descriptions, and interpretable treating gardistness and interpretable and artifications, and interpretable and artifications, and interpretable and artifications, and interpretable acceptable are protected acceptable acceptable are agants, asking a compounds are layered in asking being an artificial and are asked and are appropriated and are appropriated asking asking and are appropriated.

- 3. Semposition according to Claim 7, characterized in that it contains vitamin A, untimole sold or case of its dorivativas, or benneyl paraside, as a just for tenating acco.
- 9. Composition unsording to Claim 7, characteristed in that it contains minomidil as antiless on this requests equat and 8-cauboursethyleysteine on octopines as anytophorise agent.
- 10. Composition according to Claim 7, characterized in that is contains agreeatin or excussed as antifungal.
- 11. Composition according to Claim 7, characterized in that it contains aluminium chlorids as astringent.
- 11. Composition according to Chain 7, characterised in that it contains erythromycia or terracycline as tatilization.
- 13. Composition seconding to Chain 7, challecarized in that it contains viderables as antiviral agent.
- il. Composition acces may to Claim 7, characterism in

what it contains siculding hydrochloside as our hypernensive.

- 13. Composition according to Claim 7, characterised in these to epathine tradityeirs as maddileter.
- 13. Composition recording to their V, descentificate in them is personal a sugardade for expension in particular for expension of the personal continuity grant for expension of the particological disprists.
- 19. Composition According to Claim 7; observe the Composition as apprehension of Composition as apprehension, agrees, agrees.
- 10. Semposition recording to Claim 7, compresentated in that it contains a chromoglytate on entiallympen.
- 18. Occupatition associately to disim 7, charmotorial to them it contains a phenothiasing derivative as employmentation.
- an. Composition according to Claim 7, characterised in that it contains the opidemic growth factor (XII) or exercit factor of paperatic nature.
- 31. Desposition according to Claim 7, characterized in these it contains carifolia or theoretylline as neurostimulant.
- 12. Composition according to Claim 7, characterison in that it contains a limbium salt as antidepropagant.
- 35. Composition recording to Claim 7, characterized in Class 25 contains engrateins as instantl economic temperate ampleyed in neurobiological messanth.
- Mai Composition resording to Chain 7, that interiors in